Amendments to the Specification:

Please replace paragraph [0177] on page 19 of the application as filed, with the following amended paragraph:

[0177] The analysis was followed up for the antibody targeted against Hda1, for which it was determined the antibody was raised against a 21 amino acid peptide with the sequence TDGLNNIIEERFEEATDFILD (SEQ ID NO:22).

Comparison of this sequence with the sequence cross comparison of the 7 reactive proteins shows that the region of highest similarity (see Figure 3C) is entirely contained within the 21 amino acid sequence of the peptide (Figure 3D). To confirm this peptide as a common epitope on the proteins that cross-react with the anti-Hda1 antibody, arrays were probed with the antibody in the presence of an excess amount of the immunizing peptide; a peptide of similar length but different sequence was used as a control. As shown in Figure 2C, the Hda1 blocking peptide inhibits the interaction of anti-Hda1 with its cognate antigen Ynl021W-Hda1as well to each of the seven cross-reacting proteins. No inhibition of anti-Hda1 binding to these proteins was observed with the control peptide.

Please replace paragraph [0191] on page 21 of the application as filed, with the following amended paragraph:

[0191] Amino Acid Composition of Hda1 blocking peptide. Anti-Hda1 blocking peptide was purchased from Santa Cruz Biotechnologies, Inc (Santa Cruz, CA; 0.2ug/ul in 1XPBS). Peptide was analyzed for amino acid composition using a Beckman 7300 amino acid analyzer at the HHMI Biopolymer Keck Foundation Bioresearch Laboratory at Yale University. Comparison of the amino acid composition with the linear sequence of Hda1 was used to determine the peptide sequence: TDGLNNIIEERFEEATDFILD (SEQ ID NO:22).

Please replace paragraph [0192] on page 21 of the application as filed, with the following amended paragraph:

[0192] All antibodies were purchased from commercial vendors (See experimental protocols). For each antibody, the amount of cognate antigen present on the array by [PUT DESCRIPTION OF HOW GST WAS USED TO CALCULATE AMOUNT OF PROTEIN IN METHODS AND REFER TO IT HERE]. The antibody concentration used to probe the protein arrays was determined by titering each antibody for maximal reactivity with its cognate antigen. The number of proteins having a signal to background ratio greater than or equal to 2.0 is reported.

Please replace paragraph [0195] on page 22 of the application as filed, with the following amended paragraph:

[0195] In order to assess the utility of sequence analysis in predicting cross reactivity, all yeast proteins were searched for either the 8 amino acid epitope core sequence NNIIEERF (SEQ ID NO:1) or the 20/21 amino acid immunogenic peptide sequence TDGLNNIIEERFEEATDFILD (SEQ ID NO:22). The top matches are presented in Table 4. In addition to the observed cross-reactive proteins, a large number of proteins are identified with similarly high sequence conservation which show no empirical evidence of cross reactivity. Thus, although sequence analysis is useful in explaining the observed cross-reactivity, it is clearly insufficient to predict it.

Table 2. Sequence alignments for the 3 regions of highest homology based on a comprehensive 8 amino acid window sequence comparison. Sequence alignment is shown for 12 amino acids – the 8 amino acid core (bold in all, underlined for reference sequence YNL021W) and 2 amino acids on both N' and C'. Identities are in red.

<u>Sequence</u>	<u>Protein</u>	Identity (in 8 aa
<u>core)</u>		
Region 1		
EEENSLSTTSKS (SEQ ID NO:2)	YNL021W	
ESEESSTNSVI (SEQ ID NO:3)	YDR469W	.625
EQADSSSLTSFS (SEQ ID NO:4)	YLR332W	.5
VMENLLTTAGVS (SEQ ID NO:5)	YMR110C	.5
TDEGSYSTSIKS (SEQ ID NO:6)	YDL204W	.5

Region 2

FN epindsii sk (seq	TD NO:7)	YNL021W		
GGEPINSSVASN (SEQ		YLR332W		.625
KN EPYIDKII SK (SEQ	ID NO:9)	YDL204W		.625
FNETINKIIESK (SEQ	ID NO:10)	YMR110C		.5
MNYLIEQSNILK (SEQ	ID NO:11)	YDR469W		.375
	_			
Region 3				
GL nniieerf ee (seq	ID NO:12)	YNL021W		
AS ndiieekf yd (seq	ID NO:13)	YLR332W		.75
TINKIIEEHDTP (SEQ	ID NO:14)	YMR110C		.625
NQNVKIEESSEP (SEQ	ID NO:15)	YDR469W		.5
NLFNNEHENFDE NLFN	NRHENFDE (SEQ	ID NO:16)	YDL204W	.375

Please replace paragraph [0196] on page 22 of the application as filed, with the following amended paragraph:

[0196]

Table 3. Sequence alignment of the immunogenic peptide region with best matches from each of the 4 'cross-reactive' proteins. The 8 amino acid core from region 3 (Figure 1) is in bold for all sequences, and underlined in the reference sequence

<u>Sequence</u>	<u>Protein</u>
TDGL nniieerfe eatdfild(seq id no:17)	YNL021W
SVAS ndiieekfy deognels (seq id no:18)	YLR332W
KDFHRNKIESVLNETTKLMND (SEQ ID NO:19)	YMR110C
FHKN ynkvvekte pyidkiip <u>(seq id no:20)</u>	YDL204W
SSST nsvieesse pkisklen <u>(seq id no:21)</u>	YDR469W

Please replace paragraph [0197] on pages 22 and 23 of the application as filed, with the following amended paragraph:

[0197]

Table 4.

Table 4.		
Sequence	<u>Protein</u>	Identity
TDGLNNIIEERFEEATDFILD(SEQ ID	YNL021W	1.000
NO:22)		
TNGRNIIIEEIEASRTSFTLY (SEQ ID	YDR291W	0.476
NO:23) TDYLKNIIVENSGTSGDEDVD(SEQ ID	YIL075C	0.429
NO:24)	1110,00	0.123
RDYLNSYIEERLQEEHLDINN (SEQ ID	YKL201C	0.429
NO:25)		
KTDLVNFIEERFKTFCDEELE (SEQ ID	YKR054C	0.429
NO:26) TVLENKKIEEGKETAVDREED (SEQ ID	YKL188C	0.429
NO:27)		
IEGLNIISSGTFESLQDFVLQ(SEQ ID	YNL193W	0.429
NO:28)		
TDASNGYDEELPEEEQEFSDD(SEQ ID	YNL124W	0.429
<u>NO:29)</u>		
SYYLNCIIEENFKEMTRKLQR(SEQ ID	YNL126W	0.429
NO:30)	VDD 401C	0.381
GQFLENFLELNLNEVTDLIRD (SEQ ID NO:31)	YDR481C	0.381
TLSAGNACPGWDEDANDDILD(SEQ ID	YBR092C	0.381
No:32)		0.00=
TDIFKNCLENQFEITNLKILF (SEQ ID	YKL057C	0.381
NO:33)		
DDDDDDEDEEEEEEVTDQLED(SEQ ID	YFR033C	0.381
NO:34)	VII 1/00	0.201
VDGKGNETEEDDIKFIKGILD (SEQ ID NO:35)	YJL168C	0.381
DDGLPNGITLIGKKFTDYALL(SEQ ID	YBR208C	0.381
No:36)	1 BR200C	0.501
TISLIHEIEKIFEEDIHFEQN (SEQ ID	YHR184W	0.381
NO:37)		
FQGGLDIIKESLEEDPDFLQH(SEQ ID	YDR098C	0.381
NO: 38)	X/I D 440XI	0.201
TDYLFDYREVLESLGLDIILD (SEQ ID NO:39)	YLR443W	0.381
QFLLSKIIEARISGAFFEIWD(SEQ ID	YDL231C	0.381
NO:40)	1002310	0.501
TEFYNNYSMQVREDERDYILD(SEQ ID	YDL040	0.381

Sequence Protein Identity

NO:41)